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- Polymethylaluminoxane of enhanced solution stability.
- Delymethylaluminoxane compositions having increased solution stability in organic solvent and, in some cases, increased polymerization activity, which comprise moieties derived from an organic compound containing an electron-rich heteroatom (e.g., from Group V and/or Group VI of the Periodic Table) and a hydrocarbyl moiety. These compositions can be formed by reaction of trimethylaluminum, the organic compound, and water. They can also be formed by reaction of pre-formed polymethylaluminoxane the organic compound, which can be a hydrocarbyl group-containing compound containing an electron-rich heteroatom (e.g., oxygen, nitrogen, or sulfur), either with or without an active hydrogen atom connected to the heteroatom. The organic compound, in some embodiments, has also been found to be effective in increasing the catalytic activity of modified polymethylaluminoxane which, in addition to methyl, contain C2 and higher alkyl groups for enhanced solution stability.

BACKGROUND OF THE INVENTION

Polymethylaluminoxane, which is traditionally formed from the reaction of trimethylaluminum and a source of water, is used as a catalyst component with a metallocene catalyst component in olefin polymerizations. As indicated in European Patent Publication No. 393,358, it is difficult to form homogeneous polymethylaluminoxane solutions having good storage stability due to precipitation or gel formation in such solutions upon storage. This European patent publication mentions the reaction of polymethylaluminoxane with a C_3 to C_3 branched alkylaluminum compound in organic solvent to yield a homogeneous solution. A representative alkylaluminum compound which can be used is triisobutylaluminum.

An earlier disclosure of modified polymethylaluminoxane containing C₂ or higher alkyl groups, which has enhanced solubility in organic solvent is European Patent Publication No. 372,617 which describes compositions containing certain alkyl groups, e.g., isobutyl, n-butyl, and n-hexyl groups. This earlier European patent publication (which corresponds to U.S. Patent No. 5,041,584) does not specifically address the issue of enhancing the storage stability of conventional polymethylaluminoxane solutions which are not so modified.

A more recent U.S. patent which describes the use of tri-n-alkylaluminum compounds, having from 2 to 20 carbon atoms in the alkyl groups, to aid in the solubilization of polymethylaluminoxane in hydrocarbon solvent is U.S. Patent No. 5,066,631 to S.A. Sangokoya.

All of the above references focus upon the use of organoaluminum compounds as either reagents or additives to yield homogeneous polymethylaluminoxane solutions.

Japanese Patent Publication No. 49293/92 describes aluminoxane solutions comprising an aluminoxane, an aromatic hydrocarbon and a polar compound which can be either a halogenated hydrocarbon or a compound not containing active hydrogen but having at least one species of oxygen, sulfur, nitrogen, or phosphorus atom. O-dichlorobenzene was actually used in the Examples in this patent publication with a mention of dioxane in the preparation of the aluminoxane.

SUMMARY OF THE INVENTION

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The present invention, in one embodiment, is a novel, polymethylaluminoxane, having enhanced solution stability against precipitation or gelation during storage and, in some cases, enhanced polymerization activity. Another embodiment of the present invention relates to the improvement of the catalytic activity of certain modified polymethylaluminoxane compositions (of the type disclosed in U.S. Patent No. 5,041,584), which can already possess good solubility and solution stability due to the presence of a sufficiently high level of C₂ or higher alkyl groups, by using certain of the additives, to be described in greater detail below, which are useful in enhancing the solution stability of conventional polymethylaluminoxane.

A polymethylaluminoxane composition in accordance with this invention, is formed by the use, as either an original reagent or as a later additive, of an organic compound containing an electron-rich heteroatom and hydrocarbyl substituents to achieve the enhanced solution stability characteristics and, in some cases, enhanced activity in olefin polymerization for the polymethylaluminoxane which results. Representative compounds which contain these features include organic hydrocarbyl compounds containing electron-rich heteroatoms with active hydrogen atoms and organic hydrocarbyl compounds containing electron-rich heteroatoms without the additional presence of active hydrogen atoms. The polymethylaluminoxane composition of the present invention can be formed, for example:

- (1) by originally reacting and/or complexing trimethylaluminum, the compound containing the electronrich heteroatom and the selected hydrocarbyl moiety or moieties, and water as reagents; or
- (2) by combining the compound containing the electron-rich heteroatom and the hydrocarbyl moiety or moieties with a pre-formed polymethylaluminoxane. Modified polymethylaluminoxane of the type described in U.S. Patent No. 5,041,584, where a portion of the methyl groups in conventional polymethylaluminoxane are replaced by higher alkyl groups, is intended to be embraced by the term "polymethylaluminoxane" as used herein to the extent that the proportion and/or nature of the higher alkyl groups (i.e, the C₂ or higher alkyl groups) is not sufficient to confer the desired degree of solution stability on the entire system.

DETAILED DESCRIPTION OF THE INVENTION

Conventional, unmodified polymethylaluminoxane comprises or contains the recurring unit

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where R is methyl, as one of the principal and essential repeating units. The product may be linear, cyclic, or mixtures thereof. This material has well-recognized post-precipitation problems in organic solvent media upon storage, as previously described. It is formed by reacting trimethylaluminum with a source of water. Many specialized techniques for doing so are known to persons of ordinary skill in the art. U.S. Patent No. 5,041,585 to D. L. Deavenport et al. describes an especially preferred means for reacting trimethylaluminum with an atomized spray of water and is incorporated herein by reference, as illustrating such a preferred technique. The use of an aluminoxane reaction moderator, i.e., a preformed polymethylaluminoxane, for use in such a procedure is described in U.S. Serial No. 712,310, filed June 7, 1991, which is also incorporated herein by reference.

In accordance with the present invention, more enhanced organic solvent solubility characteristics, with a concomitant reduction of post-precipitation or gelation problems, and, in some cases, enhanced polymerization activity, can be achieved for such polymethylaluminoxane compositions by utilization of a certain type of organic compound, as either a reagent in forming the polymethylaluminoxane or as an additive to a previously formed polymethylaluminoxane. The organic compound which has been found to be useful in accordance with the present invention has two essential features. The first is at least one electron-rich heteroatom (e.g., containing one or more unshared electron pair(s)), such as one or more heteroatoms from Groups V and/or VI of the Periodic Table of the Elements, such as oxygen, nitrogen, phosphorus and/or sulfur. In many cases, the compound will also have an active hydrogen atom attached to the heteroatom as in the case of alcohols and thiols, but this is not essential since trialkyl amines and dialkyl ethers, which lack an active hydrogen atom but which have an electron-rich nitrogen or oxygen atom, are effective. The second feature is hydrocarbyl substitution which will generally be, in the aggregate with the combination of the carbon/hydrogen content of the molecule, sufficiently large to give the requisite degree of solubility for the polymethylaluminoxane. Representative hydrocarbyl groups include alkyl of, for example, C4 or higher, aryl, alkaryl, and aralkyl. In the case of certain compounds, however, such as trialkyl amines or dialkyl ethers, the hydrocarbyl groups, in the aggregate, should generally give a total carbon content for the hydrocarbyl groups in the compound of at least eight carbon atoms.

Generally, the amount of electron-rich heteroatom-containing compound to be utilized in accordance with the invention will be present at up to about 15 mole%, based on the weight of polymethylaluminoxane present or to be formed, in a preferred embodiment, although higher amounts can be used to achieve solution stability. In the latter cases, however, reduction in the catalytic activity of the treated polymethylaluminoxane is sometimes observed. Generally speaking, the person of ordinary skill in the art will probably select a level of from about 0.1 mole % to about 10 mole %, based on the amount of polymethylaluminoxane present or to be formed in order to achieve the best balance of solution stability and catalytic activity.

One way to practice the invention is to include the selected amount of the compound, containing the electron-rich heteroatom(s) and hydrocarbyl group(s), with the trimethylaluminum-containing reagent, normally employed to synthesize the polymethylaluminoxane (and, optionally, a preformed polymethylaluminoxane reaction moderator), followed by reaction of the resulting mixture with water to form a polymethylaluminoxane product. Alternatively, the compound can be placed in an appropriate hydrocarbon solvent to which the trimethylaluminum-containing reagent and water are thereafter added either simultaneously or in sequence. In both of these embodiments, the compound, which is used in accordance with the present invention, is added during the process of forming the polymethylaluminoxane and is thereby incorporated in the final solubilized polymethylaluminoxane which is formed in solubilized form in the solvent.

If desired, the compound can alternatively be added to a hydrocarbon solvent containing a pre-formed, solubilized polymethylaluminoxane with, for example, heating to achieve the desired degree of interaction and thereby achieve the results intended for the present invention.

While the precise manner in which the electron-rich heteroatom-containing, hydrocarbyl group-containing compound functions to achieve the positive effect of the present invention is not known, it is believed that such a compound may somehow either react with or coordinate with the polymethylaluminoxane or free trimethylaluminum by virtue of the electron-rich nature of the heteroatom. This coordination/reaction might prevent undesired coordination between adjacent polymethylaluminoxane chains or oligomeric/polymeric structures, which would normally occur in the absence of the compound, leading to decreased solubility of

such resulting materials in organic solvent solution. Alternatively, addition of the compound containing the electron-rich heteroatom may impart enhanced solution stability by interacting primarily with free trimethylaluminum in the solution to reduce its reactivity thereby interfering with the aging reaction between free trimethylaluminum and polymethylaluminoxane. These explanations are set forth as a possible theory of operation, and there is no intention on the part of the present inventors to be bound by such a theory merely because it is given as a potential explanation for the results observed.

As indicated earlier, it is within the purview of the present invention to use compounds containing heteroatoms, such as those from Groups V and VI of the Periodic Table of the Elements, which are capable of donating electrons (i.e., a so-called electron-rich site) to the electron deficient aluminum atoms of the polymethylaluminoxane in solution to, presumably, form complexes or reaction products which result in increased solution stability for the resulting product. One class of compound for use herein are compounds containing the electron-rich heteroatom site and at least one active hydrogen atom. Yet another class of useful compounds are those containing the previously described heteroatom which do not contain an active hydrogen atom. Non-limiting examples of such reagents containing such a hydrogen atom include compounds having the following general structures (where R is the selected hydrocarbyl group): ROH; RCO₂H; RNH₂; R₂NH; RSH; RCONH₂; RCONHR; RN(CH₂CH₂OH)₂; R₂NCH₂CH₂OH; RC(O)NHCH₂CH₂OH; RC(S)OH; and R₂PO₂H. Other compounds for use include, as representative examples, compounds containing an electron-rich heteroatom which do not contain an active hydrogen atom and which are of the following general formulae: RCONR2; RSR; ROR; RC(O)OR; RC(O)R; RC(S)R; and RC(O)H, where R is as defined above. An especially preferred R group is higher alkyl (C4 or higher) which is optionally branched at the β carbon or a carbon further removed from the a carbon atom. The preferred mode of addition is to react the active hydrogen containing hydrocarbyl moiety with preformed polymethylaluminoxane. However, incorporation into the final product can also be achieved by adding the modifier to the trimethylaluminumcontaining reagent in a suitable solvent followed by addition of water, or by adding the compound to the solvent before water and trimethylaluminum-containing reagent are added, to produce the stabilized polymethylaluminoxane/solvent composition.

Examples of such electron donor compounds, which form one type of effective compound for use herein, include the trihydrocarbyl amines, e.g., tri-n-octylamine, the trihydrocarbylphosphines, e.g., triphenylphosphine, and the dihydrocarbyl ethers, e.g., diphenylether. Ethers and amines with a total carbon content in their hydrocarbyl groups of C₈ or higher, are a preferred class of compounds to use. Two particular compounds of this general type which are particularly preferred from the standpoints of performance, availability, and cost are tridodecylamine and dibutylether.

A variety of general factors can be used to assist in the selection of particular, appropriate electron-rich heteroatom-containing and hydrocarbyl group-containing compound:

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- (1) If substituents are present on the backbone or nucleus of the selected compound, they should not give rise to an undue amount of steric hindrance in the vicinity of those portions of the compound (namely, the heteroatom or heteroatoms) which are believed to coordinate or react with the preformed aluminoxane or in forming the desired aluminoxane. Examples include: t-butanol; t-pentanol; cyclohexanol; 2,2-dimethyl-3-pentanol; 2,3-dimethyl-3-pentanol; butylated hydroxy toluene; 1,1-diphenylethanol; triethylsilanol; diphenylamine; and triphenylamine.
- (2) The use of multifunctional reagents in which the various functional groups are capable of reacting with, or in forming, more than one methylaluminoxane molecule so as to lead to bridged, or ultrahigh molecular weight species which readily precipitate are also not deemed to be preferred. Examples include: triethylene glycol monomethyl ether 1,6-hexanediol; 4-methoxyphenol; 4-nitrophenol; and dioxane.
- (3) Solution stability shows increase as the hydrocarbyl group increases in size and molecular weight. For example, methanol, when used as the alcohol reagent, yields either no stability enhancement or poor stability enhancement beyond four days, whereas dodecanol imparts stability for fourteen days under the test procedure to be described below. Alcohols of intermediate size imparts stability enhancements which are intermediate in scope: 2-ethyl-1-butanol (nine days); and 1-hexanol (seven days).
- (4) There is a concentration factor. With para-nonylphenol for example, the stability was observed to increase as the molar percent was raised, e.g., at 0.1, 0.5, 1.0, 5.0 and 10.0 mole % levels, respectively. Post-precipitation was observed within a few days with 0.1 and 0.5 mole % levels using an accelerated aging test protocol. The 1.0 mole % level lasted for several weeks before noticeable precipitation was observed. The 5.0 and 10.0 mole % concentrations showed no tendency to post-precipitate.

It is deemed that modified polymethylaluminoxane of the type shown in U.S. Patent No. 5,041,584 will not have the degree of poor storage stability experienced by conventional polymethylaluminoxane, particularly if the type and/or quantity of higher alkyl groups therein is adequate to achieve such solution stability.

Even though one embodiment of the present invention, i.e., enhanced solution stability is not as much in need of achievement with such modified compositions, it has unexpectedly been found that the catalytic activity of such a modified polymethylaluminoxane can nevertheless be increased by use of certain of the compounds described herein, e.g., the trialkylamines and the dialkyl ethers which are used with conventional polymethylaluminoxane for improved solution stability as well.

The present invention is further illustrated by the Examples which follow.

EXAMPLE 1

Procedure for Testing Stability

Candidates for use as a stabilizer in solutions of polymethylaluminoxane (PMAO) in toluene were tested using a test protocol which, for convenience, would accelerate the potential gelation or precipitation phenomenon normally observed with polymethylaluminoxane-solvent compositions over the period of many weeks to several months, depending upon the ambient temperature during storage. For example, PMAO solutions with aluminum concentrations of 10%-11% will normally show storage stabilities, before gelation/precipitation is noted, of less than about three months when stored in steel containers exposed to outside ambient temperatures in Texas. The PMAO was formed by use of the water aspiration technique described in U.S. Patent No. 5,041,585. Initial testing was conducted to verify the effectiveness of a variety of potential stabilizers. This involved addition of 1 mole % concentrations (molar ratio of additive/Al concentration in PMAO = 1:100) of the chemical to PMAO/toluene which contained 10 wt% soluble aluminum. This blend of material was transferred to a 50 ml Wheaton® vial and recapped with a Teflon® coated liner (to prevent interaction of toluene with rubber liner) under a nitrogen atmosphere in a dry box. The vial was placed in an oil bath maintained at 50-55 °C.

The vial contents were observed on a daily basis for changes in appearance and documented appropriately. Experiments were terminated after thirty days. Substrates that inhibited or substantially reduced solids formation under these conditions were subjected to additional testing. In the above tests, controls were run against a majority of the compounds tested in the same experiment. All controls showed gel or solids formation at four days or sooner.

Further experimentation was conducted to determine optimum concentrations of potential stabilizers. Varying mole % concentrations of the substrate were tested using the procedure previously described. These vials were observed on a daily basis and sampled at two week intervals to determine the aluminum content. At the end of the thirty day period the results were evaluated.

Table 1 lists the results in stability testing of substrates for PMAO/toluene. PMAO/toluene from the same lot was used for all testing. Both initial and final aluminum values were determined in most cases. The percent soluble aluminum loss after the thirty day heating period was also reported for a majority of the runs.

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5			Aluminum Loss (%)	20.0 20.4 20.4 10.2 15.3		11.0 13.0 9.0 4.0 6.1 3.1
10			Aluminum PMAO/Toluene After 30 Days (Wt %)	8.0 7.8 7.8 8.8 8.3 8.2		8.88.99.99.99.99.99.99.59.59.59.59.59.59.59.
15		oluene	ls/Gel			
20		s for PMAO/I	First Solids/Gel Formation Observed (Days)	ববববব পব		44 44 12 13 14 30 30
25		g of Substrate	Substrate (Mole %)	000000	ALCOHOLS	2331111111
35		Stability Testing of Substrates for PMAO/Toluene	Soluble Aluminum in PMAO/Toluene with Substrate (Wt %)	0.00 0.00 0.00 0.00 0.00 0.00 0.00		10.0 10.0 10.0 10.0 9.8 9.8
40						
45	TABLE 1 (p.1)		Substrate	Control Control Control Control Control Control		Methanol Ethanol Propanol Butanol Hexanol Octanol Decanol Dodecanol
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Note: Non-active hydrogen containing categories of compounds are designated by an asterisk.

	w nu		
5	Aluminum Loss (%)	9.2 17.0 10.0 14.0 10.2 9.2 10.2 5.1 20.0	3.1 3.1 11.1 19.4 20.0 20.0 16.8 40.8 35.4 3.1
10	50°C Aluminum PMAO/Toluene After 30 Days (Wt %)	8.8.8.8.8.8.8.8.8.8.8.8.8.8.8.8.8.8.8.	9.5 9.5 9.0 7.9 8.0 discontinued 7.9 5.8 6.2
15	e le		
20	Stability Testing of Substrates for PMAO/Toluene Soluble Aluminum in First Solids/Gel PMAO/Toluene With Substrate Substrate Observed (Wt %) (Mole %)	048rv00004w	30 30 26 immediate immediate 4 immediate 1
25 30	of Substrates Substrate (Mole %)		10 10 11 11 11 11 11 11 11 11 11 11 11 1
	tability Testing (Soluble Aluminum in PMAO/Toluene with Substrate		
35	ity le num Tolu Subs	9.8 10.0 10.0 9.8 9.8 9.8 9.8 9.8	9.8 9.1 9.8 9.8 9.5 9.5 9.6
	Stability Test Soluble Aluminum in PMAO/Toluene with Substra (Wt %)		o
40		ol -per -per nol	roxy T) ohol thanol 1ycol ther ol
45	TABLE 1 (p. 2) Substrate	t-Butyl alcohol t-Pentyl alcohol 2-Ethoxyethanol Cyclohexanol 2,2-Dimethyl-3-pentanol 2,3-Dimethyl-3-pentanol 2,4-Dimethyl-3-pentanol 2-Ethyl-1-butanol Benzyl alcohol	Nonylphenol Nonylphenol Nonylphenol Butylated hydroxy toluene (BHT) Phenethyl alcohol 4-Nitrophenol 1,1-Diphenylethanol Triethylene glycol monobutyl ether 4-Methoxyphenol IRGANOX 1076 brand'

octadecyl 3-(3,5-di-tert-butyl-4-hydroxy-phenyl)-propionate

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5		Aluminum Loss (%)		9.5		17.4		34.7		20.0		23.0		13.3
10	one @ 50°C	Aluminum PMAO/Toluene After 30 Days (Wt %)		8.9		8.1		6.4		8.0		1.7		8.5
15	Stability Testing of Substrates for PMAO/Toluene @ 50°C	First Solids/Gel Formation Observed (Days)						iate		5**		4**		S.
20	trates fo	First Soli Formation Observed (Days)		13	٦,	4		immediate	*S:	,	*	7	YDE	
25	ng of Subs	Substrate (Mole %)	THIOL	1	SILANOL	-	DIOL		ESTERS*		KETONE*	1	ALDEHYDE	-
30	Testi		•											
35	Stability	Soluble Aluminum in PMAO/Toluene with Substrate (Wt %)		9.8		9.8		9.8		10.0 9.5		10.0		8.6
40						loi		0]		e te				yde
45	TABLE 1 (p. 3)	Substrate		1-Octanethiol		Triethylsilanol		1,6-Hexanediol		Ethylbenzoate Dodecylacetate		Acetophenone	•	Decyl aldehyde
50														

5			Aluminum Loss (%)		1.0		15.6	8.7 18.0		5.2	2.0		10.2 3.1 1.0 12.5 3.2
10		J.05	Aluminum PMAO/Toluene After 30 Days (Wt %)		9.7 9.8		8.1	8.4		9.1	9.6		8.8 9.7 9.7 9.2
15		AO/Toluene @	First Solids/Gel Formation Observed (Days)					s		80 4 0	0.00		23801
20		es for PM/	First Soli Formation Observed (Days)	ACIDS	22 20		S	11	AMINES	71	30 30 15	AMINES	11 20 28 28 3 3
25		of Substrat	Substrate (Mole %)	PHOSPHINIC ACIDS	, <u></u>	ACIDS	1		PRIMARY AMINES	cv	1.5.	SECONDARY AMINES	
30		Stability Testing of Substrates for PMAO/Toluene	Soluble Aluminum in PMAO/Toluene with Substrate (Wt %)		9.8 9.8		9.6	9.2 10.0		9.6 9.8 8	9.8 8.8 8.8		ფ. ფ
35		Stabil	Soluble Aluminur PMAO/To with Sul (Wt %)				· [opylamine		ine mine
45	TABLE 1 (p. 4)		Substrate		Phenylphosphinic acid Diphenylphosphinic acid		Benzoic acid	z-bibenzylcarboxyllc acid Oleic acid		ylamine Ylamine	Decylamine Decylamine 3,3-Diphenylpropylamine		Dibutylamine Dioctylamine Dicyclohexylamine Diphenylamine 2,2-Dipyridylamine
50	TABI		qns		Phe Dip		Ben	z-Blbe acio Oleic		Oct Dec	Dec 3,3		90 00 00 00 00 00 00 00 00 00 00 00 00 0

5			Aluminum Loss (%)		7.3	0	2	;	6.3	٠,٠	0.0		o. o	! !	:	;	;	6.3	0.0	15.6 8.3
10		ე。09	Aluminum PMAO/Toluene After 30 Days (Wt %)		8.9	ر ب	C		0.6	υ. 		ν. c	J.	;	:	;	1	0.6	9.6	8.1 8.8
15		ne @	[a																	
20		for PMAO/Tolue	First Solids/Gel Formation Observed (Days)	VES*	**9	010 30		over 23						over 23	over 23	over 23	19	14	15	* * * 9
25 30		of Substrates	Substrate (Mole %)	TERTIARY AMINES*		-			0.1	0.3	0.5	⊣ ι	n o	0.5	0.75				۰.	, -
35		Stability Testing of Substrates for PMAO/Toluene $^{ m 6}$ $50^{ m \circ}{ m C}$	Soluble Aluminum in PMAO/Toluene with Substrate (Wt %)		9.6	y]- , ,	ກັດ	. c.	9.6	9.6	9.6	8,6	9.4	9.5	9.5	9.5	9.5	9 0	9.6	9.6 9.6
40		0,				clohexyl		•	ગુ	nine	Je	Je	ne	u ~	· a	ره ده	amine	_	niline	
45	TABLE 1 (p. 5)		Substrate		Triethylamine	iethylcy	amine Tributulamine	Tribexvlamine	Tri-n-octylamine	Tri-n-octylamin	Tri-n-octylamine	Tri-n-octylamine	Tri-n-octylamine	Tridodecylamine	Tridodecylamine	י פֿי	Trihexadecylam	(ARMEEN 316)	N.N-Diethylani	Triphenylamine Pyridine
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45	40	35	25	20	10	5
TABLE 1 (p. 6)	Stab	ility Testing	of Substrates	Stability Testing of Substrates for PMAO/Toluene @ 50°C	J.05 é	
Substrate	Sol Alu PMA Wit (Wt	Soluble Aluminum in PMAO/Toluene with Substrate (Wt %)	Substrate (Mole %)	First Solids/Gel Formation Observed (Days)	Aluminum PMAO/Toluene After 30 Days (Wt %)	Aluminum Loss (%)
ARMOSTAT 310 brand ² ARMOSTAT 410 brand ³ ARMOSTAT 710 brand ⁴	od * 3 2	0.00 6.00 6.00 6.00 6.00 6.00 6.00 6.00	ETHOXYLATED AMINES 1 1 1 1	MINES 21 over 23 over 23 over 23	1111	1111
			ETHERS*			
Dibutylether Diphenylether Furan Dioxane		8.6 9.6 9.6 9.8		29 26 6** 6**	9.6 9.6 7.8	2.0 2.0 12.5 16.4
N,N-bis(2-hydroxyethyl) N,N-bis(2-hydroxyethyl) N,N-bis(2-hydroxyethyl) predominantly saturated	xyethy xyethy xyethy atural		C16-C18 alkyl amine a 5C C8-C14 alkyl amine C6-C18 alkyl amine, cont stearyl amine.	l amine a 50/50 ratio of saturated to unsaturated groups amine, containing mainly unsaturated alkyl groups. e.	ated to unsatura turated alkyl gr	ted groups oups.

5		ne Aluminum ays Loss (%)	9.4		16.7 14.6	ن
10	50°C	Aluminum PMAO/Toluene After 30 Days (Wt %)	8.7		8.0 8.2	at 50°C/55°
15	AO/Toluene @	First Solids/Gel Formation Observed (Days)	*		* *	ys when aged
25	rates for PM	First Sol Formation e Observed (Days)	PHOSPHINE* 1 6**	SULFIDE*	3**	ss than 7 da
30	ing of Subst	Substrate (Mole %)	PHOS.	SUL	1 1	irogen atom nation in le
35	Stability Testing of Substrates for PMAO/Toluene @ 50°C	Soluble Aluminum in PMAO/Toluene with Substrate (Wt %)	9.6		9.6	an active hydds or gel forr
40	0,		Je			:king soli
45	TABLE 1 (p. 7)	Substrate	Triethylphosphine		Diethyl sulfide Thiophene	* Compounds lacking an active hydrogen atom ** results with solids or gel formation in less than 7 days when aged at 50°C/55°C.
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The Table set forth above gives data regarding the effectiveness of two of the compounds specifically mentioned as possible solution stability additives in Japanese Patent Publication No. 49293/1992 when tested in an accelerated aging test at 50 °-55 °C:

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Compound	First Solids/Gel Formation Observed (Days)
Triethylamine	6
Dioxane	6

In both cases, the solids or gel formation was first observed in six days. Hence, it is deemed unexpected that it is possible to achieve solution stabilities for conventional polymethylaluminoxane of no less than one week, often considerably longer in an accelerated aging test, using certain types of non-active hydrogen containing compounds as described herein, which, in general, contain, in the aggregate, hydrocarbyl groups of about C_8 or above. The stability data set forth in the Japanese patent publication is under ambient conditions where non-gelation of polymethylaluminoxane, as illustrated in Example 5 hereinafter, can extend for periods well over one week in the absence of any stabilizer. Accelerated aging at elevated temperature (for example, $50 ext{-}55 ext{-}C$) will generally result in solids formation or gelation in less than one week (for example four or five days) particularly if the aluminum concentration is in the range of 5%-10%. It thus appears from comparing Example 5 with the stability data discussed above that the stabilities shown in the Japanese patent publication at ambient conditions correspond to the present findings, under elevated temperature conditions, that the triethylamine and dioxane additives exert little, if any, effect on solution stability of the polymethylaluminoxane.

EXAMPLE 2

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Ethylene Polymerization Procedure

A one liter stainless steel autoclave was heated to $100\,^{\circ}$ C for one hour to remove moisture and then placed under a nitrogen purge while cooling. The autoclave was then pressurized (45 psig) with ethylene and vented twice. The autoclave was then filled with 500 cc of anhydrous, high purity toluene from Aldrich Chemical Co., was pressurized with ethylene and was heated to $80\,^{\circ}$ C. The ethylene was then vented, and the system was repressurized and vented. At this point, aluminoxane/toluene/stabilizer solution (containing 4 x 10^{-3} moles Al) was added by syringe. After about one minute of stirring, a freshly prepared solution containing 1 x 10^{-7} moles of zirconocene dichloride (Cp_2ZrCl_2) in toluene was added. The system was then pressurized to 45 psig with ethylene and the polymerization was conducted at $80\,^{\circ}$ C for fifteen minutes. The polymerization was terminated by venting the ethylene. The polymer was dried to constant weight and an activity value was calculated on the basis of gPE/(gZr.atm.hr). Table 2 shows the ethylene polymerization activities of PMAO/toluene solutions containing certain substrates, respectively:

TABLE 2

	Substrate	Mole%	Activity (10 ⁶ gPE/gZr.atm.hr)	% Gain or (% Decrease)
5	Control	0	1.66	0
	Nonylphenol	1	1.72	4
	Nonylphenol	3	1.60	(4)
	Nonyiphenol	5	1.23	(26) ·
	Nonyiphenol	10	0.64	(61)
10	Dodecanol	1	1.52	. (8)
	Dodecanol	3	1.05	(37)
	Dodecanol	5	0.88	(47)
	IRGANOX 1076 brand	1	1.45	(13)
	Decylamine	1	1.71	3
15	Decylamine	3	1.35	(19)
	Decylamine	5	0.85	(49)
	Dibutylether	1	2.12	28*
	Dioctylether	1	1.78	7
	Diphenylether	1	1.68	1
20	Dioctylsulfide	1	1.78	7
	Octylamine	1	1.58	(5)
	Dioctylamine	1	1.94	17*
	Triethylamine	1	1.53	(8)
	Diphenylphosphinic acid	1	1.18	(29)
25	Tri-n-octylamine	1	2.39	44*
	Tri-n-octylamine	5	1.93	16*
	Quinuclidine	1	1.72	4
	Quinuclidine	3	0.67	(60)
	ALIQUAT brand	1	0.19	(89)
30	Triethylphosphine	1	0.85	(49)
	Diphenylphosphinic acid	1	1.18	(29)
	ARMOSTAT 410 brand	1	1.78	7
	ARMOSTAT 710 brand	1	2.06	24*
	ARMOSTAT 1800 brand	1	1.61	(3)
35	Dibutylether**	1	1.66	0
	ARMEEN 312 brand	0.75	2.03	22*
	ARMEEN 312 brand	1	2.15	30*

^{*} particularly preferred % gains were achieved.

EXAMPLE 3

Three test substrates were added to PMAO in toluene at a molar ratio of 1:100 substrate to aluminum. Ethylene polymerization tests were conducted at several different stages of aging using the procedure described in Example 2. All aging times are at room temperature unless otherwise indicated. The identity of the tested substrate and the results of the polymerization activities are given in Table 3:

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^{**} substrate was introduced during the production of the PMAO/toluene composition.

TABLE 3

5	Aging Conditions	Polymer. Activ (10° PE/g Zr.at	ity tm.hr) % Gain or	(Decrease)
		No Subst	rate (Control)	
10	None	1.66		
		Trioctyla	amine Substrate	
15	0 Days 3 Days 5 Days 60 Days*	2.16 3.16 3.19 3.34	30 90 92 101	
20	•	Diectyla	rino Substrato	
		Dioctylai	mine Substrate	
25	O Days 9 Days at 50°C	1.58 2.45	(5) 48	
		Dioctyle	ther Substrate	
30	O Days 6 Days	1.78 2.79	7 68	
35	* 30 days ambient.	at 50°C.	followed by	30 days at

Two ethylene polymerization tests were conducted using the procedure of Example 2 except that the test conditions were 40 °C and one atmosphere ethylene. The cocatalyst used in the tests was a modified polymethyl-aluminoxane-heptane solution which is described in U.S. Patent No. 5,041,584. The difference in the two tests is the presence in the second of trioctylamine at a molar ratio of 1:100 (amine to aluminum). The following results were obtained:

TABLE 4

Amine Present (Mole %)	Polymerization Activity (10 ⁵ g PE/g Zr.atm.hr)	% Gain or (Decrease)
0	2.9	
1.0	5.4	86

EXAMPLE 5

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This Example is presented to illustrate that solutions of toluene containing polymethylaluminoxane (PMAO) and no stabilizer are subjected to solids formation and/or gelation much more quickly at ambient temperature than at elevated temperature. The samples were taken either from various stages of an actual reaction run to make PMAO ("reactor samples") or were made by dilution of the end product from such a run ("diluted samples").

PART A

Purpose: Preparation of reactor samples.

Procedure: Samples were taken at three states (25%, 50% and 75% charge of water and trimethylaluminum (TMAL)/toluene added at a concentration of 0.5 mole water per mole of aluminum) and at the conclusion of a pilot plant run to produce PMAO toluene. Each sample was analyzed for aluminum content. Preparation of the intermediate material was completed in glassware. A 1000 ml flask equipped with a stirring bar, sampling port with stopcock and septum, thermowell and nitrogen inlet/outlet was used. A nitrogen blanket was maintained on the flask throughout the procedure. A quantity of unfinished material was transferred to the flask. The apparatus was placed in an oil bath mounted on a heating/stirring plate. Stirring was initiated, and the bath temperature was lowered to 0 °C using dry ice. The bath and flask temperatures were monitored by thermocouple. Water was added (to a total of 0.75 mole of water per mole of aluminum) from a position approximately 1/2 inch above the surface of the liquid using a 5 ml syringe equipped with a Luer-Lok® stopcock and a 10 inch 22 gauge needle. Water was added at an approximate rate of 10 drops per minute (a temperature range of 0 to 5 °C was maintained by making adjustments to water addition rates). The solution was slightly cloudy with a few solid chunks following this step. The flask was heated at 80°C for sixty minutes. The flask was removed from the oil bath and cooled to room temperature. The solids were allowed to settle and the supernatant was removed and analyzed for aluminum content. The Table below contains information regarding sample preparation:

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PREPARATION OF PMAO/TOLUENE					
Stage of Sampling in Pilot Plant Run	Wt% Aluminum in Unfinished Sample	Wt of PMAO/ Toluene (g)	Vol.of Water Added (ml)	Wt% Aluminum of Final Product	
25%	4.5	352.6	1.5	3.8	
50%	6.9	383.0	3.2	6.3	
75%	9.1	371.6	4.5	8.4	
100%	10.6	371.6	0.0	10.6	

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PART B

Purpose: Preparation of diluted samples.

Procedure: Plant toluene (99.6% minimum purity) was dried overnight using molecular sieve. PMAO/toluene containing 10.6 wt% aluminum (from Procedure A) was diluted using the dried solvent to produce solutions with 4.8%, 7.0% and 8.4 wt% aluminum.

PART C

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Purpose: Determine stability of PMAO/toluene with varying aluminum concentrations at differing temperatures.

Procedure: Approximately 40 mls of each sample prepared in Procedures A and B was transferred to two 50 ml Wheaton® vials and recapped with a Teflon® coated liner (to prevent interaction of toluene with rubber of liner) in a dry box. A vial of each sample was placed in oil baths maintained at a temperature of 50 °C. The other vial was stored at room temperature (18-23 °C). The vial contents were observed on a daily basis for 47 days and intermittently thereafter for 195 days. Changes in appearance were documented appropriately. All sample vials contained solids at the end of the 195 day period. Results are listed in the following table:

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STABILITY TESTING				
Temp (°C)	Initial Conc. (% Soluble Al)	First Solids/Gel (days)		
RT*	3.8	over 47		
50*	3.8	6 days		
RT**	4.8	43 days		
50**	4.8	5 days		
RT*	6.3	28 days		
50*	6.3	5 days		
RT**	7.0	27 days		
50**	7.0	5 days		
RT*	8.4	12 days		
50*	8.4	5 days		
RT**	8.5	27 days		
50**	8.5	5 days		
RT*	10.6	12 days		
50*	10.6	5 days		

^{*} reactor sample

The percent loss of soluble aluminum rose in all cases with increased storage temperature with the loss being generally less than 10% at room temperature and 30 °C, over 10% at 40 °C, and over 20% at 50 °C.

The foregoing Examples are intended to merely illustrate certain embodiments of the present invention and, for that reason, should not be construed in a limiting sense. The scope of protection sought is set forth in the claims which follow.

Claims

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- 1. A polymethylaluminoxane composition, normally prone to precipitation from organic solvent over time, which is dissolved in organic solvent and which shows increased solution storage stability, said polymethylaluminoxane comprising moieties derived from an organic compound containing at least one electron-rich heteroatom and at least one hydrocarbyl group selected from the group consisting of compounds containing an active hydrogen atom and compounds which contain no active hydrogen atom, the presence of such moieties from the organic compound being effective to enhance the solvent storage stability of the composition comprising the dissolved polymethylaluminoxane and solvent, with the proviso that when the compound contains no active hydrogen atom there is no solids or gel formation for at least seven days when aged at a temperature of about 50 °C to about 55 °C.
- A composition as claimed in Claim 1 wherein the heteroatom is selected from the group consisting of Group V and Group VI of the Periodic Table of the Elements.
- 3. A composition as claimed in Claim 1 wherein the heteroatom is selected from the group consisting of oxygen, nitrogen, phosphorus, and sulfur.
 - 4. A composition as claimed in Claims 1-3 wherein the organic compound comprises an electron-rich heteroatom and an active hydrogen atom.
- 5. A composition as claimed in Claims 1-3 wherein the organic compound comprises an electron-rich heteroatom and no active hydrogen atom.
 - 6. A composition as claimed in Claims 1-3 wherein the organic compound is an ether.
 - 7. A composition as claimed in Claim 1 wherein the organic compound is an amine.
 - 8. A composition as claimed in Claim 6 wherein the organic compound is dibutyl ether.
 - 9. A composition as claimed in Claim 7 wherein the organic compound is tridodecyl amine.

^{**} dilution sample

- 10. A modified polymethylaluminoxane composition, which comprises, in addition to methyl groups, hydrocarbyl groups higher in molecular weight than methyl for enhanced solvent solubility, which modified polymethylaluminoxane is normally stabilized against precipitation from organic solvent over time and which is dissolved in organic solvent, said polymethylaluminoxane comprising moieties derived from an organic compound containing at least one electron-rich heteroatom and at least one hydrocarbyl group, the presence of such moieties from that organic compound being effective to enhance the catalytic activity of the modified polymethylaluminoxane composition.
- 11. A composition as claimed in Claim 10 wherein the heteroatom is selected from the group consisting of Group V and Group VI of the Periodic Table of the Elements.
 - 12. A composition as claimed in Claim 10 wherein the heteroatom is selected from the group consisting of oxygen, nitrogen, phosphorus, and sulfur.
- 13. A composition as claimed in Claims 10-12 wherein the organic compound comprises an electron-rich heteroatom and an active hydrogen atom.
 - 14. A composition as claimed in Claims 10-12 wherein the organic compound comprises an electron-rich heteroatom and no active hydrogen atom.
 - 15. A composition as claimed in Claim 10 wherein organic compound is an ether.
 - 16. A composition as claimed in Claim 10 wherein the organic compound is an amine.
- 25 17. A composition as claimed in Claim 15 wherein the organic compound is dibutyl ether.
 - 18. A composition as claimed in Claim 16 wherein the organic compound is tridodecyl amine.

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